

Claim 58 (Cancelled)

Claim 59 (Currently Amended): The transgenic ~~mammal~~ rat of claim 57, wherein the MMP-13 comprises the sequence of SEQ ID NO: 1 or SEQ ID NO: 21.

Claim 60 (Currently Amended): The transgenic ~~mammal~~ rat of claim 55, wherein the repressor-activator fusion polypeptide is a chimeric tetracycline repressor-VP16 transcription activator polypeptide and the regulatable promoter is a Tn10 sequence linked to a portion of the CMV IE promoter.

Claim 61 (Currently Amended): The transgenic ~~mammal~~ rat of claim 60, wherein the regulatable promoter comprises the sequence of SEQ ID NO: 2.

Claim 62 (Currently Amended): The transgenic ~~mammal~~ rat of claim 55, wherein the Type II collagen degradation results in a loss of proteoglycan, cleavage of Type II collagen into a TCA degradation product, a change in joint function, joint space narrowing, destruction of cartilage, a change in growth plate morphology, fibrillation and loss of articular cartilage, osteophyte formation, or combinations thereof.

Claim 63 (Currently Amended): The transgenic ~~mammal~~ rat of claim 55, wherein the ~~joint~~ chondrocyte-specific promoter is a Type II collagen promoter.

Claim 64 (Currently Amended): A transgenic rat whose genome comprises:

(a) a nucleotide sequence encoding a constitutively enzymatically active human matrix metalloproteinase that cleaves Type II collagen, wherein the nucleotide sequence encoding the metalloproteinase is operatively linked to a tetracycline-regulatable promoter; and

(b) a nucleotide sequence encoding a repressor-activator fusion polypeptide that binds to the tetracycline regulatable promoter in the absence of tetracycline or a tetracycline analog and does not bind to the regulatable promoter in the presence of tetracycline or tetracycline analog, which nucleotide sequence encoding the repressor-activator fusion polypeptide is operatively linked to a ~~joint~~ chondrocyte-specific promoter, wherein expression of the metalloproteinase is capable of being repressed in the rat until adulthood, and wherein the metalloproteinase is capable of being expressed in the rat during adulthood to a level sufficient to cause Type II collagen degradation in the joints of the rat.

Claim 65 (Currently Amended): The transgenic rat of claim 64, wherein the matrix metalloproteinase is constitutively enzymatically active MMP-13, the tetracycline-regulatable promoter is tet07, the repressor-activator fusion polypeptide is tTA, and the ~~joint~~ chondrocyte-specific promoter is a Type II collagen promoter.

Claim 66 (Previously presented): The transgenic rat of claim 64, wherein the Type II collagen degradation results in a loss of proteoglycan, cleavage of type II collagen into a TCA degradation product, a change in joint function, joint space narrowing, destruction of cartilage, a change in growth plate morphology, fibrillation and loss of articular cartilage, osteophyte formation, or combinations thereof.

Claim 67 (Currently Amended): A method for producing degradation of Type II collagen in the joints of a transgenic ~~non-human-mammal~~ rat, which method comprises:

- (a) maintaining the transgenic ~~mammal~~ rat of claim 55 in presence of the transcription activator protein-binding compound until adulthood; and
- (b) activating expression of the matrix metalloproteinase in the transgenic ~~mammal~~ rat by withholding the compound from the ~~mammal~~ rat after the ~~mammal~~ rat has

Claim 75 (Currently Amended): A transgenic non-human ~~mammal~~ rat whose genome comprises:

(a) a nucleotide sequence encoding a constitutively enzymatically active human matrix metalloproteinase that cleaves Type II collagen, wherein the nucleotide sequence encoding the metalloproteinase is operatively linked to a regulatable promoter; and

(b) a nucleotide sequence encoding a transcription activator protein that binds to the regulatable promoter in the presence of a transcription activator protein-binding compound and does not bind to the regulatable promoter in the absence of the compound, which nucleotide sequence encoding the transcription activator protein is operatively linked to a chondrocyte-specific promoter;

wherein expression of the metalloproteinase is capable of being repressed in the ~~mammal~~ rat until adulthood, and wherein the metalloproteinase is capable of being expressed in the ~~mammal~~ rat during adulthood to a level sufficient to cause Type II collagen degradation in the joints of the ~~mammal~~ rat.

Claim 76 (Currently Amended): The transgenic ~~mammal~~ rat of claim 75, wherein the matrix metalloproteinase is selected from the group consisting of MMP-1, MMP-8, and MMP-13.

Claim 77 (Currently Amended): The transgenic ~~mammal~~ rat of claim 76, wherein the matrix metalloproteinase is MMP-13.

Claim 78 (Cancelled)

Claim 79 (Currently Amended): The transgenic ~~mammal~~ rat of claim 77, wherein the MMP-13 comprises the sequence of SEQ ID NO: 1 or SEQ ID NO: 21.

Claim 80 (Currently Amended): The transgenic ~~mammal~~ rat of claim 75, wherein the chondrocyte-specific promoter is a Type II collagen promoter.

Claim 81 (Currently Amended): The transgenic ~~mammal~~ rat of claim 75, wherein the transcription activator protein is a chimeric polypeptide comprising a transactivator domain linked to an ecdysone receptor ligand-binding domain, and wherein the transgenic ~~mammal~~ rat further comprises a nucleotide sequence encoding a retinoid X receptor (RXR), which nucleotide sequence encoding RXR is operatively linked to a chondrocyte-specific promoter.

Claim 82 (Currently Amended): The transgenic ~~mammal~~ rat of claim 75, wherein the transcription activator protein is a chimeric polypeptide comprising a transactivator domain linked to a progesterone receptor ligand-binding domain.

Claim 83 (Currently Amended): The transgenic ~~mammal~~ rat of claim 75, wherein the transcription activator protein is a chimeric polypeptide comprising a transactivator domain linked to a steroid binding domain.

Claim 84 (Currently Amended): The transgenic ~~mammal~~ rat of claim 75, wherein the Type II collagen degradation results in a loss of proteoglycan, cleavage of type II collagen into a TCA degradation product, a change in joint function, joint space narrowing, destruction of cartilage, a change in growth plate morphology, fibrillation and loss of articular cartilage, osteophyte formation, or combinations thereof.

Claim 85 (Currently Amended): A method for producing degradation of Type II collagen in the joints of a transgenic non-human ~~mammal~~ rat, which method comprises:

- (a) maintaining the transgenic ~~mammal~~ rat of claim 75 in the absence of the transcription activator protein-binding compound until adulthood; and
- (b) activating expression of the matrix metalloproteinase in the transgenic ~~mammal~~ rat by administering the compound to the ~~mammal~~ rat after the ~~mammal~~ rat has

reached adulthood such that the matrix metalloproteinase degrades Type II collagen in the joints of the ~~mammal~~ rat.

Claim 86 (Currently Amended): A method for producing degradation of Type II collagen in the joints of a transgenic ~~non-human mammal~~ rat, which method comprises:

- (a) maintaining the transgenic ~~mammal~~ rat of claim 81 in the absence of ecdysone, an ecdysone analog, or dexamethasone until adulthood; and
- (b) activating expression of the matrix metalloproteinase in the transgenic ~~mammal~~ rat by administering ecdysone, an ecdysone analog, or dexamethasone to the ~~mammal~~ rat after the ~~mammal~~ rat has reached adulthood such that the matrix metalloproteinase degrades Type II collagen in the joints of the ~~mammal~~ rat.

Claim 87 (Currently Amended): A method for producing degradation of Type II collagen in the joints of a transgenic ~~non-human mammal~~ rat, which method comprises:

- (a) maintaining the transgenic ~~mammal~~ rat of claim 82 in the absence of mifepristone (RU 486) until adulthood; and
- (b) activating expression of the matrix metalloproteinase in the transgenic ~~mammal~~ rat by administering mifepristone (RU 486) to the ~~mammal~~ rat after the ~~mammal~~ rat has reached adulthood such that the matrix metalloproteinase degrades Type II collagen in the joints of the ~~mammal~~ rat.

Claim 88 (Previously presented): The method according to claim 86, wherein the Type II collagen degradation results in a loss of proteoglycan, cleavage of type II collagen into a TCA degradation product, a change in joint function, joint space narrowing, destruction of cartilage, a change in growth plate morphology, fibrillation and loss of articular cartilage, osteophyte formation, or combinations thereof.

Claim 89 (Previously presented): The method according to claim 87, wherein the Type II collagen degradation results in a loss of proteoglycan, cleavage of type II collagen into a TCA degradation product, a change in joint function, joint space narrowing, destruction of cartilage, a change in growth plate morphology, fibrillation and loss of articular cartilage, osteophyte formation, or combinations thereof.

Claim 90 (Currently Amended): A method for evaluating the potential of a composition to counteract degradation of Type II collagen in joints of a transgenic ~~non-human mammal~~ rat, which degradation results in a phenotypic change selected from the group consisting of loss of proteoglycan, cleavage of Type II collagen into a TCA degradation product, a change in joint function, joint space narrowing, destruction of cartilage, a change in growth plate morphology, fibrillation and loss of articular cartilage, osteophyte formation, and combinations thereof, which method comprises:

(a) providing a first and second transgenic ~~non-human mammal~~ rat of claim 55

(b) ~~in which a phenotypic change has been produced by activation of~~ activating expression of the metalloproteinase at the same age during adulthood of the transgenic ~~non-human mammals~~ rats, wherein expression of the metalloproteinase results in a phenotypic change selected from the group consisting of loss of proteoglycan, cleavage of Type II collagen into a TCA degradation product, a change in joint function, joint space narrowing, destruction of cartilage, a change in growth plate morphology, fibrillation and loss of articular cartilage, osteophyte formation, and combinations thereof;

(b) (c) administering the composition to the first transgenic ~~non-human mammal~~ rat; and

(e) (d) comparing the phenotype of the first transgenic ~~non-human mammal~~ rat to which the composition was administered with the phenotype of the second transgenic ~~non-human mammal~~ rat in which the composition was not administered,

wherein any less extensive development in the nature or extent of the ~~phenotypic change~~ phenotype in the first transgenic ~~non-human mammal~~ rat or any increased

Claim 93 (Currently Amended): A method for evaluating the potential of a composition to counteract degradation of Type II collagen in joints of a transgenic ~~non-human mammal~~ rat, which degradation results in a phenotypic change selected from the group consisting of loss of proteoglycan, cleavage of Type II collagen into a TCA degradation product, a change in joint function, joint space narrowing, destruction of cartilage, a change in growth plate morphology, fibrillation and loss of articular cartilage, osteophyte formation, and combinations thereof, which method comprises:

(a) providing a first and second transgenic ~~non-human mammal~~ rat of claim 75;

(b) ~~in which a phenotypic change has been produced by activation of~~ activating expression of the metalloproteinase at the same age during adulthood of the transgenic ~~non-human mammals~~ rats, wherein expression of the metalloproteinase results in a phenotypic change selected from the group consisting of loss of proteoglycan, cleavage of Type II collagen into a TCA degradation product, a change in joint function, joint space narrowing, destruction of cartilage, a change in growth plate morphology, fibrillation and loss of articular cartilage, osteophyte formation, and combinations thereof;

(b) (c) administering the composition to the first transgenic ~~non-human mammal~~ rat; and

(e) (d) comparing the phenotype of the first transgenic ~~non-human mammal~~ rat to which the composition was administered with the phenotype of the second transgenic ~~non-human mammal~~ rat in which the composition was not administered,

wherein any less extensive development in the nature or extent of the ~~phenotypic change~~ phenotype in the first transgenic ~~non-human mammal~~ rat or any increased length of time required for the ~~phenotypic change~~ phenotype to develop in the first transgenic ~~non-human mammal~~ rat that has been administered the composition relative to the ~~phenotypic change~~ phenotype in the second transgenic ~~non-human mammal~~ rat, indicates the potential of the composition to counteract the phenotypic change.

Claim 94 (Currently Amended): A method for evaluating the potential of a composition to counteract degradation of Type II collagen in joints of a transgenic ~~non-human mammal~~ rat, which degradation results in a phenotype selected from the group consisting of loss of proteoglycan, cleavage of Type II collagen into a TCA degradation product, a change in joint function, joint space narrowing, destruction of cartilage, a change in growth plate morphology, fibrillation and loss of articular cartilage, osteophyte formation, and combinations thereof, which method comprises:

(a) providing a first and second transgenic ~~non-human mammal~~ rat of claim 81;

(b) ~~in which a phenotypic change has been produced by activation of~~ activating expression of the metalloproteinase at the same age during adulthood of the transgenic ~~non-human mammals~~ rats, wherein expression of the metalloproteinase results in a phenotypic change selected from the group consisting of loss of proteoglycan, cleavage of Type II collagen into a TCA degradation product, a change in joint function, joint space narrowing, destruction of cartilage, a change in growth plate morphology, fibrillation and loss of articular cartilage, osteophyte formation, and combinations thereof;

(b) (c) administering the composition to the first transgenic ~~non-human mammal~~ rat; and

(e) (d) comparing the phenotype of the first transgenic ~~non-human mammal~~ rat to which the composition was administered with the phenotype of the second transgenic ~~non-human mammal~~ rat in which the composition was not administered,

wherein any less extensive development in the nature or extent of the ~~phenotypic change~~ phenotype in the first transgenic ~~non-human mammal~~ rat or any increased length of time required for the ~~phenotypic change~~ phenotype to develop in the first transgenic ~~non-human mammal~~ rat that has been administered the composition relative to the ~~phenotypic change~~ phenotype in the second transgenic ~~non-human mammal~~ rat, indicates the potential of the composition to counteract the phenotypic change.

Claim 96 (Currently Amended): A method for evaluating the potential of a composition to counteract degradation of Type II collagen in joints of a transgenic ~~non-human mammal~~ rat, which degradation results in a phenotypic change selected from the group consisting of loss of proteoglycan, cleavage of Type II collagen into a TCA degradation product, a change in joint function, joint space narrowing, destruction of cartilage, a change in growth plate morphology, fibrillation and loss of articular cartilage, osteophyte formation, and combinations thereof, which method comprises:

(a) providing a first and second transgenic ~~non-human mammal~~ rat of claim 83;

(b) ~~in which a phenotypic change has been produced by activation of~~ activating expression of the metalloproteinase at the same age during adulthood of the ~~non-human mammals~~ rats, wherein expression of the metalloproteinase results in a phenotypic change selected from the group consisting of loss of proteoglycan, cleavage of Type II collagen into a TCA degradation product, a change in joint function, joint space narrowing, destruction of cartilage, a change in growth plate morphology, fibrillation and loss of articular cartilage, osteophyte formation, and combinations thereof;

(b) (c) administering the composition to the first transgenic ~~non-human mammal~~ rat; and

(e) (d) comparing the phenotype of the first transgenic ~~non-human mammal~~ rat to which the composition was administered with the phenotype of the second transgenic ~~non-human mammal~~ rat in which the composition was not administered,

wherein any less extensive development in the nature or extent of the ~~phenotypic change~~ phenotype in the first transgenic ~~non-human mammal~~ rat or any increased length of time required for the ~~phenotypic change~~ phenotype to develop in the first transgenic ~~non-human mammal~~ rat that has been administered the composition relative to the ~~phenotypic change~~ phenotype in the second transgenic ~~non-human mammal~~ rat, indicates the potential of the composition to counteract the phenotypic change.

Claim 97 (Currently Amended): A method for producing degradation of Type II collagen in the joints of a transgenic ~~non-human-mammal~~ rat, which method comprises:

- (a) maintaining the transgenic ~~non-human-mammal~~ rat of claim 83 in the absence of mifepristone (RU 486) until adulthood; and
- (b) activating expression of the matrix metalloproteinase in the transgenic ~~mammal~~ rat by administering mifepristone (RU 486) to the ~~mammal~~ rat after the ~~mammal~~ rat has reached adulthood such that the matrix metalloproteinase degrades Type II collagen in the joints of the ~~mammal~~ rat.

Claim 98 (New) A method for evaluating the potential of a composition to counteract degradation of Type II collagen in joints of a transgenic mouse, which degradation results in a phenotypic change selected from the group consisting of loss of proteoglycan, cleavage of Type II collagen into a TCA degradation product, a change in joint function, joint space narrowing, destruction of cartilage, a change in growth plate morphology, fibrillation and loss of articular cartilage, osteophyte formation, and combinations thereof, which method comprises:

- (a) providing a first and second transgenic mouse, whose genomes each comprise:
 - (i) a nucleotide sequence encoding a constitutively enzymatically active human matrix metalloproteinase that cleaves Type II collagen, wherein the nucleotide sequence encoding the metalloproteinase is operatively linked to a regulatable promoter; and
 - (ii) a nucleotide sequence encoding a repressor-activator fusion polypeptide that binds to the regulatable promoter in the absence of a repressor-activator fusion polypeptide-binding compound and does not bind to the regulatable promoter in the presence of the compound, which nucleotide sequence encoding the repressor-activator fusion polypeptide is operatively linked to a chondrocyte-specific promoter,wherein expression of the metalloproteinase is capable of being repressed in the mouse until adulthood, and wherein the metalloproteinase is capable of being

expressed in the mouse during adulthood to a level sufficient to cause Type II collagen degradation in the joints of the mouse;

(b) activating expression of the metalloproteinase at the same age during adulthood of the transgenic mice, wherein expression of the metalloproteinase results in a phenotypic change selected from the group consisting of loss of proteoglycan, cleavage of Type II collagen into a TCA degradation product, a change in joint function, joint space narrowing, destruction of cartilage, a change in growth plate morphology, fibrillation and loss of articular cartilage, osteophyte formation, and combinations thereof;

(c) administering the composition to the first transgenic mouse; and

(d) comparing the phenotype of the first transgenic mouse to which the composition was administered with the phenotype of the second transgenic mouse in which the composition was not administered,

wherein any less extensive development in the nature or extent of the phenotype in the first transgenic mouse or any increased length of time required for the phenotype to develop in the first transgenic mouse that has been administered the composition relative to the mouse phenotype in the second transgenic mouse, indicates the potential of the composition to counteract the phenotypic change.

99 (New): The method of claim 98, wherein the repressor-activator fusion polypeptide of the transgenic mouse is a chimeric tetracycline repressor-VP16 transcription activator polypeptide and the regulatable promoter is a Tn10 sequence linked to a portion of the CMV IE promoter.

100 (New): The method of claim 98, wherein the regulatable promoter is a tetracycline-regulatable promoter and the repressor-activator fusion polypeptide-binding compound is tetracycline or a tetracycline analog.